# 中華民國專利 報 (19)(12)

(11)公告級號:341533

〔41〕中華民國87年(1998)10月01日

(51) Int + C | 6:805D1/22 805B5/03 第 P010518 號 9 引証附件

(54)名

们:在斜板上途復鋅之設備與方法

(21)中 請 案 號:86115993

(22)中諸日期:中華民國86年(1997)10月28日

〔72〕發 明 人:

金相忠

韓國

韓國

超雷夏 🔝

却國

(71)中 請 人:

浦項綜合製鐵股份有限公司

如 國

(74)代 理 人:蔡坤財 先生

1

1

## [57]申請專利範圍:

- 1.一種將鋅連續地塗覆於鋼板上的設備, 其包含:
  - 一鋅塗覆室,用以形成一鋅粉的流化床,用以將一加熱的鋼板通過該鋅粉的流化床,及用以在鋼板通過該流化床時 5. 使得鋅粉熔黏於該鋼板上:
  - 一流化床形成室,其藉由噴出一氣體來 讓鋅粉懸浮而形成鋅粉的一流化床;
  - 一旋流器,其在將鋅粉從該鋅塗覆室中 回收之後將鋅粉從氣體中分離出來以排 出該氣體,並將被分離出來的鋅粉送回 到該流化床形成室中:
  - 一轉向器,其在鋼板進入到該鋅塗覆室 中之後將鋼板的前進方向偏移:
  - 一張力滾子,其將一塗覆了鋅的鋼板的 前進方向偏移:

該鋅塗覆室包括:一從該鋅塗覆室的一 側壁連接至該流化床之粉末入口管用以 將鋅粉噴入該鋅塗覆室中:一氣體入口 管用以形成該鋅粉的一亂流及用以防止 鋅粉的滲漏:及一回收管用以重新使用 未被塗覆的鋅粉;

該氣體入口管是位在該粉末入口管之上,及該回收管是位在該粉末入口管的 底下:

該回收管連接於該鋅塗覆室與該旋流器 之間,及一抽吸泵接於該旋流器上:

- 一分隔板設在該鋅塗覆室中用以讓沒有 被塗覆的鋅粉能平順地流到該回收管 中,及用以防止鋅粉在通過該回收管之 後進入到該鋅塗覆室中;及
- 一穩定滾子·其位在該分隔板的底下。
- 2.如申請專利範圍第1項所述之設備,其中上述之加熱機構為一退火爐。
- 15. 3.如申請專利範圍第1或2項中所述之設備,其中一再加熱爐被裝設於該鋅塗覆室與該張力滾子之間。
  - 4.一種將鋅連續地塗覆於鋼板上的設備· 其包含:
- 20. 一鋅塗覆室,用以讓鋅粉熔黏於經加熱

4

10.

## 的鋼板上以形成一塗覆層:

- 一流化床形成室,其藉由噴出一氣體來 讓鋅粉懸浮而形成鋅粉的一流化床:
- 一旋流器,其在將鋅粉從該鋅塗覆室中 回收之後將鋅粉從氣體中分離出來以排 出該氣體,並將被分離出來的鋅粉送回 到該流化床形成室中:
- 一轉向器滾子,其在鋼板進入到該鋅塗 覆室中之後將鋼板的前進方向偏移;
- 一張力滾子,其將一塗覆了鋅的鋼板的 前進方向偏移:

該鋅塗覆室包括一粉末噴出管,其從該 鋅塗覆室的一側壁連接至該流化床用以 將鋅粉噴入該鋅塗覆室中;

該鋅塗覆室更包括一連接至該旋流器之 回收管用以回收未被塗複的鋅粉:及 一或多個電極,其係設置在該鋅塗覆室 中用以將鋅粉靜電地帶電,該等電極係 連接至一高電壓產生裝置。

- 5.如申請專利範圍第4項所述之設備,其 中一再加熱爐被裝設於該鋅塗覆室與該 張力滾子之間。
- 6.如申請專利範圍第4或5項所述之設備,其中該粉末噴出裝置包括:一攜粉管,其連接至該流化床形成室:一注入泵,其連接至該攜粉管:及一粉末噴出噴嘴,用來將鋅粉從注入泵噴入到該鋅塗覆室中。
- 7.如申請專利範圍第4或5項所述之設備,其中該電極是以一種與該鋅塗覆室隔離開來及一對尖的電極設在鋅塗覆室的側壁上的狀態被裝設於該鋅塗覆室中。
- 8.如申請專利範圍第6項所述之設備,其 中該電極包含一對網式電極,其是以一 種與該對尖的電極及該鋅塗覆室隔離開 來的狀態被裝設於該鋅塗覆室中。
- 9.如申請專利範圍第4項所述之設備,其 中一冷卻裝置被裝設於該鋅塗複室的底 下。

- 10.如申請專利範圍第9項所述之設備, 其中該冷卻裝置包括:一噴水噴嘴:及 一噴氣噴嘴用以在該噴水噴嘴之上形成 一空氣簾幕。
- 11.如申請專利範圍第5項所述之設備, 其中該冷卻裝置是置於該再加熱爐的底下。
- 12.如申請專利範圍第11項所述之設備, 其中該冷卻裝置包括:一噴水噴嘴:及 10. 一噴氣噴嘴用以在該噴水噴嘴之上形成 一空氣簾幕。
  - 13.如申請專利範圍第10或12項所述之設備,其中一清洗裝置被裝設於該噴水噴 嘴底下,用以清洗該鋼板的表面。
- 15. 14.一種藉由將鋼板通過一鋅塗覆室而連續地將鋅塗覆於鋼板上的方法,其包括的步驟為:

提供該鋅塗覆室一流化床之經流化的鋅粉,及經由該鋅塗覆室的一例壁將一鈍 20. 氣或還原氣體噴入該鋅塗覆室中以在該 鋅塗覆室中形成一流化床:

> 將一經加熱的鋼板(被加熱至 430-730 ℃)通過在該鋅塗覆室中之流化床用以 將鋅粉熔黏於該鋼板上以形成一塗覆 層:

- 25. 層: 在 420-650℃的溫度下重新加熱該黏附 了鋅粉的鋼板 1-20 秒鐘用以讓剩餘未 塗覆之鋅粉熔黏於該鋼板的表面上以形 成一塗覆層:及
- 30. 用一旋流器從該鋅塗覆室的底部將剩餘 未塗覆之鋅粉與一氣體一起排出,將鋅 粉與該氣體分開來以排出該氣體及將分 離出來的鋅粉送回到一流化床形成室 中。
- 35. 15.如申請專利範圍第14項所述之方法, 其中該鋅粉具有一45 μ m 的平均粒子 大小。
  - 16.如申請專利範圍第14或15項所述之方法,其中鋅粉包含了0.1-0.7重量%的。

40.

10.

- 17.如申請專利範圍第14或15項所述之方法,其中所使用的再加熱溫度為420-500℃及再加熱的時間為1-5秒鐘。
- 18.如申請專利範圍第16項所述之方法, 其中所使用的再加熱溫度為420-500℃ 及再加熱的時間為1-5秒鐘。
- 19.如申請專利範圍第14或15項所述之方法,其中所使用的再加熱溫度為500-650℃及再加熱的時間為10-20秒鐘。
- 20.如申請專利範圍第 16項所述之方法· 其中所使用的再加熱溫度為 500-650℃ 及再加熱的時間為 10-20 秒鐘。
- 21.如申請專利範圍第14或15項所述之方法,其中該鋅塗覆室使用一溫度為250 ℃或更低的環境的氣體。
- 22.如申請專利範圍第16項所述之方法, 其中該鋅塗覆室使用一溫度為250℃或 更低的環境的氣體。
- 23.一種藉由將鋼板通過一鋅塗覆室而連 續地將鋅塗覆於鋼板上的方法,其包括 的步驟為:

從一粉末供應裝置接受鋅粉,並藉由一 從底部吹出之氣體之助將鋅粉於一流化 床形成室中加以流化;

將該經流化的鋅粉從該流化床形成室經 由一注入裝置而噴入一鋅塗覆室中以在 該鋅塗覆室中形成一流化床:

將該流化床之鋅粉帶以正電荷或負電 荷:

將一鋼板加熱至 420-730℃ 並將該鋼板接地,並將該帶電荷的鋼板通過該流化床用以讓鋅粉熔黏於該鋼板上:

將在該鋅塗覆室底部上之剩餘的鋅粉與 一氣體一起送至一提流器用以將鋅粉與 該氣體分開來,及排出該氣體,並將分 離出來的鋅粉送回到該粉末供應裝置。

- 24 如申請專利範圍第 14 項所述之方法, 其中該鋅粉具有一  $45~\mu$  m 的平均粒子 大小。
- 25.如申請專利範圍第23或24項所述之方

- 法,其中鋅粉包含了 0.1-0.7 重量%的 鋁。
- 26.如申請專利範圍第23項中任一項所述 之方法,其中在鋅塗覆該鋼板之後,該 5. 鋼板於500-650℃的溫度下被再加熱, 用以將鬆弛地附著之未被塗覆的鋅粉熔 黏於該鋼板上。
  - 27.如申請專利範圍第 26 項所述之方法, 其中所使用的再加熱溫度為 420-500℃ 及再加熱的時間為 1-5 秒鐘。
  - 28.如申請專利範圍第26項所述之方法, 其中所使用的再加熱溫度為420-500℃ 及再加熱的時間為1-5秒鐘。
- 29.如申請專利範圍第23或24項所述之方 15. 法·其中該鋼板被接地,及該電極被供 應-1至-100KV或1至100KV的電壓。
  - 30.如申請專利範圍第25項所述之方法, 其中該鋼板被接地,及該電極被供應-1 至-100KV或1至100KV的電壓。
- 20. 31.如申請專利範圍第 26 、 27 或 28 項所 述之方法,其中該鋼板被接地,及該電 極被供應 -1 至 -100KV或 1 至 100KV的 電壓。
- 32.如申請專利範圍第23或24項所述之方 25. 法,其中該鋅塗覆室使用的氛圍氣體的 溫度為正常溫度至250℃。
  - 33.如申請專利範圍第25項所述之方法, 其中該鋅塗覆室使用的氛圍氣體的溫度 為正常溫度至250℃。
- 30. 34.如申請專利範圍第 26 、 27 或 28 項所 述之方法,其中該鋅塗覆室使用的氛圍 氣體的溫度為正常溫度至 250℃。
  - 35.如申請專利範圍第23或24項所述之方法,其中在該鋼板於該鋅塗覆室中鋅塗 覆之後,其被一由一噴嘴噴出的水所冷卻。
    - 36.如申請專利範圍第25項所述之方法, 其中在該鋼板於該鋅塗覆室中鋅塗覆之 後,其被一由一噴嘴噴出的水所冷卻。
- 40. 37.如申請專利範圍第 26 、 27 或 28 項所

35.

8

3 (E)

述之方法,其中在該鋼板於該鋅塗覆室 中鋅塗覆之後,其被一由一噴嘴噴出的 水所冷卻。

- 38.如申請專利範圍第23或34項中任一項 所述之方法,其中在該鋼板於該鋅塗覆 室中鋅塗覆之後,其被水所清洗。
- 39.如申請專利範圍第25項所述之方法, 其中在該鋼板於該鋅塗覆室中鋅塗覆之 後,其被水所清洗。
- 40.如申請專利範圍第 26 、 27 或 38 項所 述之方法,其中在該鋼板於該鋅塗覆室 中鋅塗覆之後,其被水所清洗。

#### 圖式簡單說明:

第一圖為一傳統鋅塗覆設備的示意圖:

第二圖為另一傳統鋅塗覆設備的示 意圖:

第三圖顯示依據本發明之鋅塗覆設 備的一實施例:

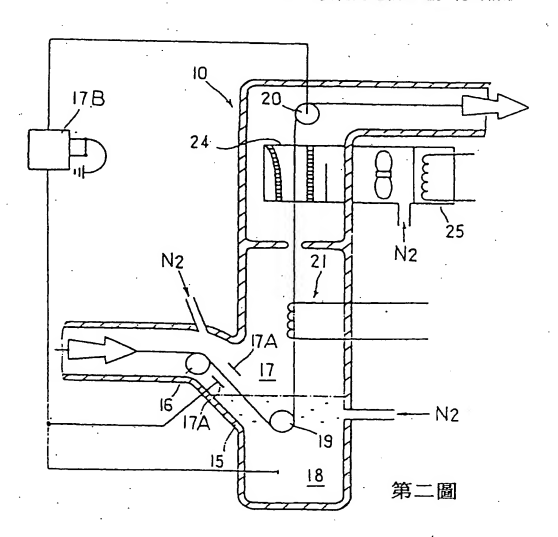
 第四圖顯示依據本發明之鋅塗覆設 備的另一實施例:

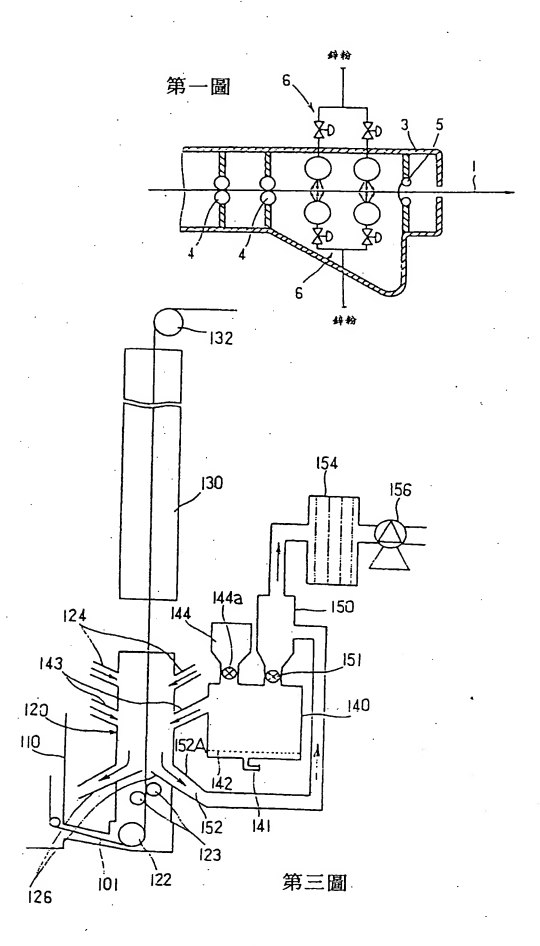
第五圖為第四圖的 A 部分的詳細圖式:

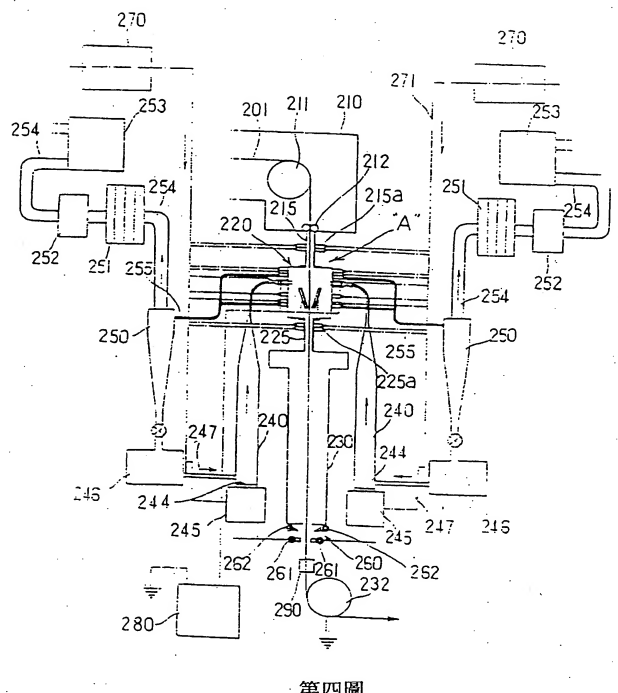
第六圖為一圖表,其顯示鋅粉末之 10. 結塊的變化對在該鋅塗覆室中之流化床的 溫度的情形:

> 第七圖為一圖表,其顯示塗覆量的 變化對氣流速率的情形:及

第八圖為一圖表,其顯示塗覆量的 15. 變化對電極電壓變化的情形。

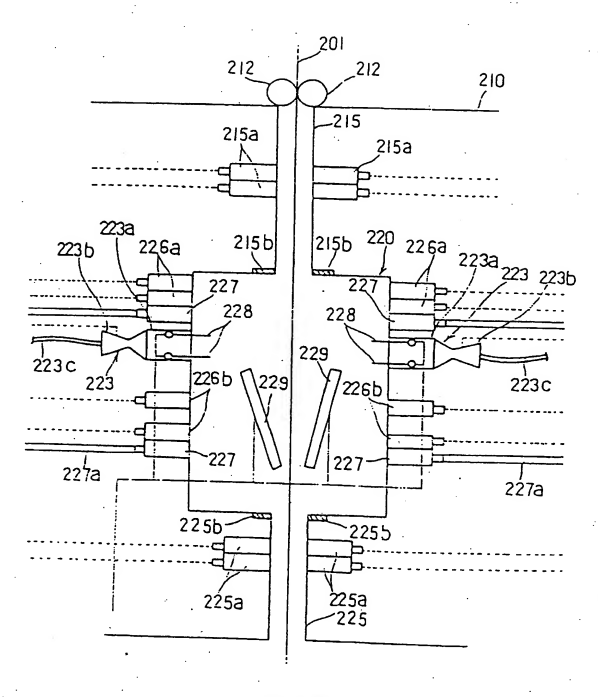




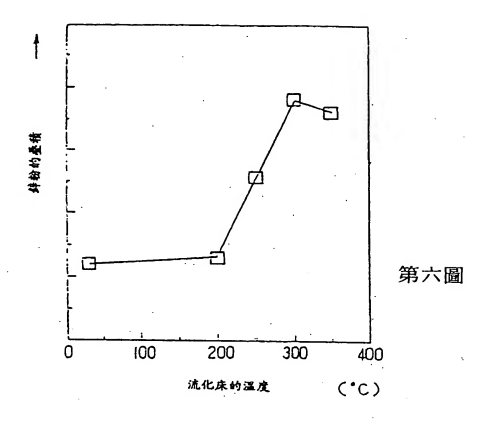


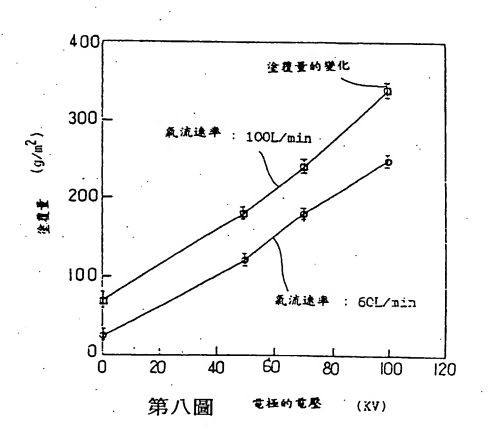
第四圖

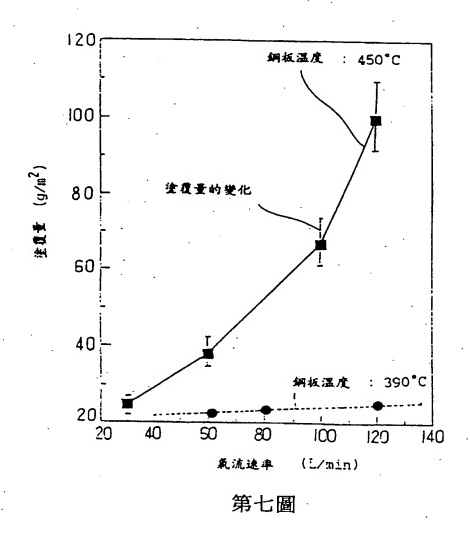
**:** 



第五圖







341533

APPARATUS FOR COATING ZINC ON STEEL SHEET, AND METHOD THEREFOR

#### BACKGROUND OF THE INVENTION

#### 1. Field of the invention

The present invention relates to an apparatus and a method for coating zinc on steel sheets for use on automobiles and electronic apparatuses. More specifically, the present invention relates to an apparatus and a method for coating zinc on steel sheets by using zinc powders.

## 2. Description of the prior art

Zinc performs an sacrificing action for steel to extend the life expectancy of steel, and therefore, conventionally zinc has been used in zinc-coating a steel sheet. There are many kinds of zinc coating methods such as hot dip galvanization, electroplating, and zinc powder-using zinc coating. coated steel sheets are mostly used as automobile body sheets and outer and inner sheets of electronic apparatuses. They are manufactured by electroplating or hot dip galvanization. The reason is as follows. That is, when a steel strip is continuously coated, not only the product quality but also the productivity and the workability have to be considered. In this respect, electroplating and hot dip galvanization are advantageous.

Electroplating is carried out in the following manner. That is, cold rolled steel sheets are made to undergo a batch annealing or a continuous annealing so as to improve mechanical

properties. Then, electroplating is carried out within an electrolyte containing zinc ions, thereby obtaining a zinc deposition layer of the target thickness. In this method, the mechanical properties which have been obtained through heat treatments are not degraded during the plating process. Further, the coated amount (deposition thickness) is varied in accordance with the applied electric power, and therefore, the coated amount can be accurately controlled.

However, it has the following disadvantages. That is, as the coated amount is increased, so much more electric power is required. Consequently, the productivity is aggravated in the case of a thick plating. Therefore, electroplating is adopted in the case where the coated amount is  $40~\text{g/m}^2$  or less for one face of steel sheets. Meanwhile, the plating speed is limited by the limit current density, and therefore, if the productivity is to be improved, the zinc coating chamber has to be long. This becomes a facility burden. Because of such limiting factors, electroplating should be preferably carried out at a strip velocity of about 200 m/min and at a coated amount of  $40~\text{g/m}^2$  or less for one face of the steel sheet.

Meanwhile, hot dip galvanization is carried out in the same facility as that of the annealing. Therefore the manufacturing cost becomes lower, and a thick zinc coating is possible. However, it has the following disadvantages. That is, a sink roll and a guide roll which are immersed in a hot dipping pot are corroded, and therefore, they have to be replaced periodically. Further, as the line speed becomes

fast, so much the resistivity of the melted zinc is increased. Therefore, the sink roll cannot move synchronously with the steel sheet to produce slips, and therefore, the surface of the steel sheet may be scratched so much as to lead to a product defect. Further, as the line speed is increased, or as the coated thickness is decreased, splashes are increased during an air wiping, with the result that the generation of dross is increased. Besides, if the zinc adhered on the surface is to be solidified, some cooling period is required, and therefore, the velocity of the steel sheet is limited to about 200 m/min. Further, the adjustment of the coated thickness is difficult, and therefore, the manufacturing becomes difficult if the coated amount per face is less than 40 g/m².

Meanwhile, a method for coating zinc by using zinc powders is disclosed in Japanese Patent Application Lid-open No. Hei-5-311388.

This method is illustrated in FIG. 1. As shown in this drawing, an object to be coated (steel sheet) 1 is heated to above the melting point of the powder metal, and zinc powders loaded on a gas are spouted by means of a spouting nozzle 6 onto the object 1 within a zinc coating chamber 3 containing a reducing atmosphere. Thus the zinc powders are melt-adhered on the steel sheet 1, thereby zinc-coating the steel sheet.

In FIG. 1, reference codes 4 and 5 indicate sealing devices.

In this method, a reducing atmosphere is used, and therefore, a flux does not have to be used. Further, compared

with the hot dip galvanization, the air wiping and the management of the melt composition are not required, and the dross generation does not occur.

However, in the case of Japanese Patent Application Laidopen No. Hei-5-311388, the zinc powders from the powder storage
chamber are not screened but directly spouted into the zinc
coating chamber. Therefore, large particles and coarse
secondary particles can adhere on the steel sheet, with the
result that the coated layer becomes irregular.

Meanwhile, another zinc coating method is disclosed in which an object to be coated is heated to 775°F (413°C) - 820°F (438°C), and zinc powders or a zinc melt is spouted, so that zinc would be coated on steel sheets (CA 866153 (7113).

In this method, however, a flux is spouted together with the zinc powders to prevent the oxidation of the steel sheet. Further, in order to improve the adherence of zinc, electrostatic charges of opposite polarities are provided on the zinc powders and the object to be coated. In this method, owing to the mechanical spouting force and the electrostatic attractions, a large coated amount can be easily obtained. Further, it can be applied to a complicated steel structure, but when it is applied to a continuous zinc coating of a steel strip, the following problems occur. (1) The high voltage electrostatic charges of opposite polarities are dangerous to workers. (2) There are necessarily loosely adhered zinc particles after the zinc coating, and these particles adhere on various rolls to cause defects called "dent". (3) Zinc

powders are released into the external air to aggravate the working environment.

Further, there are other disclosures in which electrostatic charges are utilized in coating zinc powders (U.S. Patents 5,384,165 and 5,551,981).

In these methods, zinc powders are made to adhere on steel sheets by utilizing electrostatic charges, and then, the steel sheets are heated to convert the adhered zinc powders to a coated layer.

The apparatus for these methods is illustrated in FIG. 2. As shown in this drawing, the apparatus includes a fluidized bed 18 of zinc powders, and a cooling device 24 and a heating device 21 disposed above the fluidized bed 18. A steel sheet immersed in fluidized bed 18 is made to shift its advancing direction upward to be heated by the heating device 21, so that the zinc powders would be melted. The melted zinc powders are reflowed, and then are cooled.

In FIG. 2, reference code 10 indicates a housing, 16 indicates a strip bending roll, 17 indicates a fall space, 17A indicates a plate as a part of an electrostatic charge circuit, 17B indicates a controller, 20 indicates a top deflector roll.

In these U.S. patents, steel sheets can be coated without much modifying the existing melting facility, but have the following disadvantages.

(1) When the electrostatically charged metal powders contact with the steel sheet, the surface charges are transferred to the steel sheet so as to be grounded and to

disappear. Therefore, the attractive force of the electrostatic charges which is to act as the adhering force between the steel sheet and the zinc powder is dissipated. Therefore, the zinc powders depart from the surface of the steel sheet, and therefore, there is a limit in the increase of the coated thickness.

- (2) The roller is immersed in the fluidized bed of zinc powders, and therefore, when the steel sheets moves, the zinc powders intrude into between the roller and the steel sheet, that the zinc powders may adhere on the Particularly, zinc powders speedily undergo sintering reactions above 250°C. Therefore, the zinc powders which have intruded into between the steel sheet and the roller undergo a sintering reaction owing to the latent heat of the steel sheet. coarse particles may be formed, and the dent result, phenomenon becomes more serious.
- (3) The fine zinc particles of 5-15  $\mu$ m which are used in the above patents are not well fluidized, but agglomerations occur. Therefore, the zinc particles of the fluidized bed are liable to be irregular, and therefore, if the steel sheet is put into the fluidized bed, a uniform coated layer cannot be obtained.

Further, if a reflowing is carried out after the adherence of the zinc powders, a volume contraction occurs as in the case of the powder metallurgy. Therefore, the coated layer may look as if the steel sheets has cracked. Further, if the reflowing is imperfectly carried out, the residual zinc powders of the

surface will adhere on the roller so as to form a dent defect.

### SUMMARY OF THE INVENTION

In order to solve the above described disadvantages of the conventional techniques, the present inventors carried out researches and studies, and based on the result of the researches and studies, the present inventors proposes the present invention.

Therefore it is an object of the present invention to provide a zinc coating apparatus and a method therefor, in which a fluidized bed forming chamber is provided to fluidize zinc powders in carrying out a zinc coating on a heated steel sheet, so that not only a uniformly coated layer but also a thick coated layer can be obtained.

It is another object of the present invention to provide a zinc coating apparatus and a method therefor, in which a fluidized bed forming chamber is provided to fluidize zinc powders, and the fluidized zinc powders are electrostatically charged to coat zinc on a heated steel sheet, so that not only a uniformly coated layer but also a thick coated layer can be obtained, and that an aesthetically superior coated layer can be obtained.

In achieving the above objects, the apparatus for continuously coating zinc on a steel sheet according to the present invention includes:

a zinc coating chamber for forming a fluidized bed of zinc powders, for passing a heated steel sheet through the fluidized

bed of the zinc powders, and for making the zinc powders meltadhere on the steel sheet during the passing of the steel sheet through the fluidized bed;

a fluidized bed forming chamber for forming a fluidized bed of the zinc powders by making the zinc powders suspended by spouting a gas;

a cyclone for separating the zinc powders from the gas after recovery of them from the zinc coating chamber, to discharge the gas, and to return the separated zinc powders to the fluidized bed forming chamber;

a deflector for shifting the advancing direction of the steel sheet after its admittance into the zinc coating chamber;

a tension roll for shifting the advancing direction of a zinc coated steel sheet;

the zinc coating chamber including: a powder inlet tube connected from a side wall of the zinc coating chamber to the fluidized bed to inject the zinc powders into the zinc coating chamber; a gas inlet tube for forming a turbulent flow of the zinc powders and for preventing a leakage of the zinc powders; and a recovering tube for reusing uncoated zinc powders;

the gas inlet tube being disposed above the powder inlet tube, and the recovering tube being disposed below the powder inlet tube;

the recovering tube being connected between the zinc coating chamber and the cyclone, and a suction pump being connected to the cyclone;

a separating plate provided within the zinc coating

chamber, for making the uncoated zinc powders smoothly flow to the recovering tube, and for preventing the zinc powders from flowing into the zinc coating chamber after passing through the recovering tube; and

a stabilizing roll disposed below the separating plate.

In another aspect of the present invention, the apparatus for continuously coating zinc on a steel sheet according to the present invention includes:

a zinc coating chamber for making zinc powders melt-adhere on a heated steel sheet to form a coated layer;

a fluidized bed forming chamber for forming a fluidized bed of the zinc powders by making the zinc powders suspended by spouting a gas;

a cyclone for separating the zinc powders from the gas after recovery of them from the zinc coating chamber, to discharge the gas, and to return the separated zinc powders to the fluidized bed forming chamber;

a deflector roll for shifting the advancing direction of the steel sheet after its admittance into the zinc coating chamber;

a tension roll for shifting the advancing direction of a zinc coated steel sheet;

the zinc coating chamber including a powder spouting tube connected from a side wall of the zinc coating chamber to the fluidized bed to spout the zinc powders into the zinc coating chamber;

the zinc coating chamber further including a recovering

tube connected to the cyclone, for recovering uncoated zinc powders; and

one or more of electrodes provided in the zinc coating chamber, for electrostatically charging the zinc powders, the electrodes being connected to a high voltage generating device.

In still another aspect of the present invention, the method for continuously coating zinc on a steel sheet by making the steel sheet pass through a zinc coating chamber according to the present invention, includes the steps of:

supplying fluidized zinc powders of a fluidized bed to the zinc coating chamber, and injecting an inert gas or a reducing gas into the zinc coating chamber through a side wall of the zinc coating chamber to form a fluidized bed within the zinc coating chamber;

making a heated steel sheet (heated to 420 - 730°C) through the fluidized bed within the zinc coating chamber to melt-attach the zinc powders on the steel sheet so as to form a coated layer;

reheating the zinc powder-adhered steel sheet at a temperature of 420 - 650°C for 1 - 20 seconds to make residual uncoated zinc powders melt-adhered on the surface of the steel sheet so as to form a coated layer; and

discharging residual uncoated zinc powders from a bottom portion of the zinc coating chamber together with a gas by a cyclone, to separate the zinc powders from the gas so as to discharge the gas and so as to return the separated zinc powders to a fluidized bed forming chamber.

In still another aspect of the present invention, the method for continuously coating zinc on a steel sheet by making the steel sheet pass through a zinc coating chamber according to the present invention, includes the steps of:

receiving zinc powders from a powder supply device, and fluidizing the zinc powders within a fluidized bed forming chamber by a help of a gas blown through a lower portion;

injecting the fluidized zinc powders from the fluidized bed forming chamber to a zinc coating chamber by means of an injecting device to form a fluidized bed within the zinc coating chamber;

charging the zinc powders of the fluidized bed positively or negatively;

heating a steel sheet to 420 - 730°C and grounding the steel sheet, and making the charged steel sheet pass through the fluidized bed to make the zinc powders melt-adhere on the steel sheet;

sending a residual zinc powders of a bottom portion of the zinc coating chamber to a cyclone together with a gas to separate the zinc powders from the gas, so as to discharge the gas, and so as to return the separated zinc powders to a powder supply device.

The above method further includes the step of: forming a coated layer by melt-coating the zinc powders on the steel sheet, and then, carrying out a reheating at a temperature of 420 - 650°C for 1 - 20 seconds to make residual uncoated zinc powders melt-adhere on the steel sheet.

#### BRIEF DESCRIPTION OF THE DRAWINGS

The above objects and other advantages of the present invention will become more apparent by describing in detail the preferred embodiment of the present invention with reference to the attached drawings in which:

- FIG. 1 is a schematic illustration of a conventional zinc coating apparatus;
- FIG. 2 is a schematic illustration of another conventional zinc coating apparatus;
- FIG. 3 illustrates an embodiment of the zinc coating apparatus according to the present invention;
- FIG. 4 illustrates another embodiment of the zinc coating apparatus according to the present invention;
- FIG. 5 is a detailed illustration of a portion A of FIG. 4;
- FIG. 6 is a graphical illustration showing the variation of agglomeration of the zinc powders versus the temperature of the fluidized bed within the zinc coating chamber;
- FIG. 7 is a graphical illustration showing the variation of the coated amount versus the variation of the gas flow rate; and
- FIG. 8 is a graphical illustration showing the variation of the coated amount versus the variation of the voltage of the electrode.

## DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

FIG. 3 illustrates a first embodiment of the zinc coating

apparatus according to the present invention.

As shown in this drawing, the apparatus includes:

a zinc coating chamber 120 for forming a fluidized bed of zinc powders, for passing a heated steel sheet (steel strip) 101 through the fluidized bed of the zinc powders, and for making the zinc powders melt-adhere on the steel sheet during the passing of the steel sheet through the fluidized bed;

a fluidized bed forming chamber 140 for forming a fluidized bed of the zinc powders by making the zinc powders suspended by spouting a gas;

a cyclone 150 for separating the zinc powders from the gas after recovery of them from the zinc coating chamber 120, to discharge the gas, and to return the separated zinc powders to the fluidized bed forming chamber 140;

a deflector 122 for shifting the advancing direction of the steel sheet after its admittance into the zinc coating chamber 120; and

a tension roll 132 for shifting the advancing direction of a zinc coated steel sheet.

A means for heating the steel sheet may be an annealing furnace 110 as shown in FIG. 3.

The zinc coating chamber includes a powder spouting tube 143 connected from a side wall of the zinc coating chamber 120 to the fluidized bed forming chamber 140 to inject the fluidized zinc powders into the zinc coating chamber 120.

Further, on a side wall of the zinc coating chamber 120, there is provided a gas inlet tube 124 for forming a turbulent

flow of the zinc powders and for preventing a leakage of the zinc powders. The zinc powders are supplied from a powder supply tube 143, and the gas inlet tube 124 is disposed above the powder supply tube 143.

On a side wall of the zinc coating chamber 120, there is further connected a recovering tube 152 for sending uncoated and descending zinc powders and the gas to the cyclone 150.

The recovering tube 152 should preferably include an inclined portion 152A, and this inclined portion 152A should be constituted such that it can facilitate the recovery of the zinc powders.

The zinc coating chamber 120 includes a separating plate 126. This separating plate 126 makes the uncoated zinc powders flow smoothly to the recovering tube 152, and prevents the zinc powders from flowing through a recovering tube connecting portion to the lower portion of the zinc coating chamber 120.

Beneath the separating plate 126, there is disposed a stabilizing roll 123.

The upper portion of the fluidized bed forming chamber 140 is connected to a hopper 144 which supplies the zinc powders. To the bottom of the fluidized bed forming chamber 140, there is connected a gas supply tube 141 which is connected to a gas supply source (not shown in the drawings). Below the fluidized bed forming chamber 140, there can be disposed a porous gas dispersing plate 142 which disperses the gas from the gas supply tube 141 to obtain a uniform fluidized bed.

The fluidized bed forming chamber 140 and the cyclone 150

can be provided in the number of one or more. In the present invention, even if a single fluidized bed forming chamber 140 is installed, a plurality of powder inputting tubes 143 can be connected to it, so that the zinc powders can be inputted into the zinc coating chamber 120 from a plurality of points.

The cyclone 150 is connected to a suction pump 156, and the suction pump 156 sucks the uncoated zinc powders and the gas from the zinc coating chamber 120.

A filter 154 should be preferably installed between the cyclone 150 and the suction pump 156, so that the zinc powders remaining in the gas after the gas-powder separation by the cyclone 150 can be captured.

The uncoated zinc powders remaining on the coated surface of the steel sheet should be converted into a zinc coated layer. For this purpose, the coated steel sheet is heated. In order to carry out the heating, a reheating furnace 130 should be preferably installed between the zinc coating chamber 120 and the tension roll 132.

Valves 144a and 151 are installed respectively under the hopper 144 and the cyclone 150.

In another embodiment of the present invention, the apparatus for continuously coating zinc on a steel sheet according to the present invention as shown in FIG. 4 includes:

a zinc coating chamber 220 for making zinc powders meltadhere on a heated steel sheet (strip) 201 to form a coated layer;

a fluidized bed forming chamber 240 for forming a fluidized

bed of the zinc powders by making the zinc powders suspended by spouting a gas;

a cyclone 250 for separating the zinc powders from the gas after recovery of them from the zinc coating chamber 220, to discharge the gas, and to return the separated zinc powders to the fluidized bed forming chamber 240;

a deflector 211 for shifting the advancing direction of the steel sheet after its admittance into the zinc coating chamber 220; and

a tension roll 232 for shifting the advancing direction of a zinc coated steel sheet.

The means for heating the steel sheet may consist of an annealing furnace 210 as shown in FIG. 4.

The zinc coating chamber 220 includes a powder spouting device 223 connected from a side wall of the zinc coating chamber 220 to the fluidized bed forming chamber 240 to spout the zinc powders into the zinc coating chamber 220.

The powder spouting device 223 should preferably include: a powder carrying tube 223c connected to the fluidized bed forming chamber 240; an injection pump 223b connected to the powder carrying tube 223c; and a powder spouting nozzle 223a for spouting the zinc powders from the injection pump 223b into the zinc coating chamber 220.

The power spouting device 223 is installed in the number of two on the opposite side walls of the zinc coating chamber 220.

Further, the zinc coating chamber 220 should be provided

with one or more of electrodes for electrostatically charging the zinc powders.

As shown in FIG. 5, a pair of sharp electrodes 228 should be preferably provided on the sides walls of the zinc coating chamber 220 oppositely facingly. More preferably, a pair of the sharp electrodes 228 should be provided on the side walls of the zinc coating chamber 220, and at the same time, a pair of net type electrodes 229 should be provided mutually facingly across the advancing steel sheet within the zinc coating chamber 220.

These electrodes are connected to a high voltage generator 280.

These electrodes are for electrostatically charging the zinc powders with negative or positive charges.

A recovering tube 227 which is connected to the cyclone 250 is also connected to a lower portion of the side wall of the zinc coating chamber 220, for recovering the uncoated zinc powders. The recovering tube 227 can be connected not only to the zinc coating chamber 220 but also to an upper portion of the side wall of the powder spouting device 223.

An upper sealing chamber 215 is installed between the annealing furnace 210 and the zinc coating chamber 220, while a lower sealing chamber 225 is installed between the zinc coating chamber 220 and the reheating furnace 230.

The upper sealing chamber 215 communicates to the zinc coating chamber 220. The isolation between the annealing furnace 210 and the zinc coating chamber 220 should preferably

be done by a pair of sealing rolls 212 which prevent lateral oscillations of the steel sheet, electrically ground the steel sheet, and seal the atmosphere of the annealing furnace 210.

A pair or more of gas spouting nozzles 215a should be preferably provided on the upper sealing chamber 215, so that the zinc powders can be prevented from floating up to the sealing roll 212, and that the internal pressure of the zinc coating chamber 220 can be adjusted.

The lower sealing chamber 225 should be preferably provided with a pair or more of gas spouting nozzles 225a to inject an atmospheric gas so as to adjust the internal pressure of the zinc coating chamber 220.

The upper sealing chamber 215 and the zinc coating chamber 220 should be insulated from each other by means of an insulator 215b. The lower sealing chamber 225 and the zinc coating chamber 220 should be insulated from each other by means of an insulator 225b.

If the insulators are provided as described above, then the walls of the zinc coating chamber are charged with the same polarity as that of the zinc powders, and therefore, the zinc powders can be prevented from being adhered on the walls of the zinc coating chamber.

Within the zinc coating chamber, the flow of the zinc powders should be prevented from being lamella within the internal atmosphere. For this purpose, a pair or more of gas spouting nozzles 226a should be preferably installed on the upper portion of the side wall of the zinc coating chamber 220.

Further, a pair or more of gas spouting nozzles 226b should be preferably installed on the side wall of the zinc coating chamber 220 between the powder spouting nozzle 223 and the powder recovering tube 227.

As shown in FIG. 4, a gas discharge tube 254 is connected to the top of the cyclone 250, and the gas discharge tube 254 is connected to a suction pump 253 which transfers the gas and the uncoated zinc powders from the zinc coating chamber 220 to the cyclone 250.

The gas discharge tube 254 includes a back filter 251 and a dust collector 252, and the back filter 251 filters the discharge gas, while the dust collector 252 collects fine zinc powders after their passing through the back filter 251.

Meanwhile, the bottom of the cyclone 250 communicates to the powder supply device 246, and therefore, the zinc powders which have been separated from the gas are carried to the powder supply device 246.

The lower side wall of the fluidized bed forming chamber 240 is connected to the powder supply device 246 through the powder supply tube 247, so that the zinc powders can be supplied to the fluidized bed forming chamber 240. A gas supply part 245 is connected to the bottom of the fluidized bed forming chamber 240, for supplying the fluidizing gas to the fluidized bed forming chamber 240.

A porous gas dispersing plate 244 should be preferably installed in the lower portion of the fluidized bed forming chamber 240, so that the gas supplied from the gas supply part

245 can be uniformly dispersed.

The fluidized bed forming chamber 240 and the cyclone 250 can be provided in the number of one or more.

In the present invention, even in the case where a single fluidized bed forming chamber 240 is installed, a plurality of the powder spouting device 223 can be connected to the fluidized bed forming chamber 240, so that the zinc powders can be spouting from a plurality of points into the zinc coating chamber 220.

A reheating furnace 230 should be preferably installed between the zinc coating chamber 220 and the tension roll 232, so that the coated steel sheet can be reheated, and that the residual zinc powders can be converted into a coated layer.

Meanwhile, in the case where the reheating furnace 230 is installed beneath the zinc coating chamber 220, a cooling device 260 can be installed beneath the reheating furnace 230. This cooling device 260 should preferably consist of an air spouting nozzle 262 for forming an air curtain above a water spouting nozzle 261.

A washing device 290 having a brush for washing the surface of the steel sheet can be installed beneath the water spouting nozzle 261 of the cooling device 260. In the case where this washing device 290 is installed, the residual zinc powders are removed from the surface of the steel sheet, and therefore, the workability is improved.

further, in the case where the reheating furnace 230 is installed, a holding chamber can be installed, so that the

temperature of the steel sheet under the reheating furnace can be maintained at  $500 - 650\,^{\circ}\text{C}$  to subject the coated layer to an alloying treatment.

Meanwhile, in the present invention, a gas heating device 270 should be preferably installed to heat the supplied gas to a certain temperature level.

Now the operation of the zinc coating apparatus of the present invention will be described referring to FIGs. 4 and 5.

the steel sheet 201 is heat-treated in the annealing furnace 210, and the temperature of the steel sheet is adjusted to 420 - 730°C. This steel sheet enters into the zinc coating chamber 220 by the help of the deflector roll 211 and the tension roll 232. Meanwhile, the zinc powders are supplied from the powder supply device 246 through the powder supply tube 247 into the fluidized bed forming chamber 240 to be suspended there. Then the zinc powders are transferred through the powder carrying tube 223c, the injection pump 223b and the powder spouting nozzle 223a into the zinc coating Then the zinc powders are first charged by an chamber 220. electrode 228 which is disposed near the powder spouting nozzle 223a so as to adhere on the steel sheet. Owing to a nitrogen gas or nitrogen-hydrogen gas mixture which is spouted through the gas spouting nozzle 226b of the zinc coating chamber 220, layered flows are prevented. Further, the zinc powders are completely charged by the net type electrode 229, result that the adhering efficiency is improved. Under this condition, the gas and the residual uncoated zinc powders are

sucked into the recovering tube 227 so as to double the turbulent flow effect. In order to improve the efficiency of the flow of the zinc powders, the pressure and the flow rate of the fluidized carrier gas and the auxiliary gas of the ventury tube are properly adjusted. The coated amount during the zinc coating is adjusted by the supplied amount of the zinc powders, by the amount of the gas spouted into the fluidized bed and the ventury tube, and by the voltage supplied to the electrode.

The residual zinc powders after the zinc coating are discharged through the recovering tube 227 and the gas discharge. tube 254 to the outside of the zinc coating chamber 220 by the suction pump 253. Then they are transferred to the cyclone 250, and are separated into the gas and the zinc powders. separated zinc powders are reinputted through a valve into the powder supply device, while the separated gas is discharged through the back filter 251 and the dust collector 252. order to prevent the zinc powders from being mixed into the annealing furnace 210, a nitrogen or nitrogen-hydrogen mixture gas is spouted through the gas spouting nozzle 215a of the upper sealing chamber 215 so as to form a gas curtain, and to adjust the internal pressure of the zinc coating chamber 220. Further, the oscillations of the steel sheet have to be prevented, steel sheet has to be grounded, and the atmospheres of the annealing furnace 210 and the zinc coating chamber 220 have to be isolated from each other. For these purposes, the sealing roll 212 is driven. In the lower sealing chamber 225 of the

zinc coating chamber 220 also, a nitrogen gas or nitrogenhydrogen mixture gas is spouted through the gas spouting nozzle 225a so as to form a gas curtain, and to adjust the internal pressure of the zinc coating chamber 220. The coated steel sheet is heated by the reheating furnace 230 to make the imperfectly adhered zinc powders melt-adhere on the steel sheet. Further, if necessary, the coated steel sheet is subjected to a zinc-iron alloying reaction within a holding chamber. cooling the reheated steel sheet, if a water cooling method is adopted, the cooling efficiency is improved. The steam which is generated during the water cooling is not fed into the zinc coating chamber but discharged to the outside by the help of the air curtain of the cooling device. Depending on cases, coated steel sheet can be washed before it is contacted with the tension roll, thereby completely removing the loosely adhered zinc powders. The nitrogen or nitrogen-hydrogen mixture gas is heated by the gas heating device 270 before being supplied through the gas supply tube so as to be used in the zinc coating.

Now the method for coating zinc on the steel sheet according to the present invention will be described in detail.

In the present invention, a steel sheet which has been heated to a proper temperature is contacted with solid phase metallic zinc powders which are suspended. Thus owing to the latent heat of the steel sheet, the zinc powders adhere on the surface of the steel sheet perfectly or imperfectly. Therefore, a coated layer is formed, and in the case where the adherence

is imperfect, a reheating is carried out, so that a perfect adherence can be realized through a melt-adherence.

Specifically, in the present invention, the desirable conditions for forming a perfect coated layer are as follows.

- 1) Oxides should not exist on the surfaces of the steel sheet, so that the coated layer would closely adhere on the steel sheet.
- 2) The steel sheet has to have a latent heat enough to ensure a perfect melt-adherence of the zinc powders on the steel sheet.

In a state with the Items 1 and 2 satisfied, if the following conditions are met, then a satisfactory quality in the zinc coated steel sheet can be obtained.

- 3) In order to obtain a uniform coated layer, the particles of the zinc powders should have less than a certain size.
- 4) The adherence of the coated layer should be superior, the Item 1 has to be met, and an excessive formation of an alloy layer (F phase) on the boundary between the coated layer and the steel sheet should be inhibited.
- 5) In the zinc powder adhered layer, imperfect adherences can easily occur, and these imperfectly adhered powders should be removed or made to melt-adhere for ensuring the quality of the zinc coated sheet.

In the present invention, the conditions for meeting the above Items 1-5 are as follows.

If the Item 1 is to be met, the atmosphere which is used

during the heating of the steel sheet has to be a reducing gas or a non-oxidizing gas. In the steel manufacturing industry, this condition can be satisfied in the continuous annealing furnace which is used in manufacturing cold-rolled steel sheets. Generally, the used gases are mixtures of nitrogen plus hydrogen or nitrogen plus carbon monoxide. In the general continuous annealing furnace, the formation of oxides rarely occurs, and therefore, the Item 1 can be sufficiently satisfied.

If the Item 2 is to be met, the temperature of the steel sheet should be preferably limited to 420 - 730°C, and the reason is as follows. That is, at a temperature of 419°C which is the melting point of zinc, there can occur imperfect adherences or adherences through diffusion reactions. However, if a sound coated layer is to be ensured, the reheating step is necessary.

At the reheating step, when the zinc powders melt-adhere, the external appearance of the steel sheet may be aggravated due to the volume contraction. Further, too much load is imposed on the reheating furnace. On the other hand, if the steel sheet is heated to above 730°C, the mechanical properties may be aggravated, and a zinc-iron alloying reaction is excessively promoted, with the result that the adherence of the coated layer is adversely affected.

If the Item 3 is to be met, the average particle size of the zinc powders should be preferably limited to less than 45  $\mu m$  (-325 mesh). In the case where the coated amount is as low

as 50 g/m<sup>2</sup>, if the average particle size is more than 45  $\mu$ m, then the adhered powder amount is too small, with the result that some parts of the steel sheet may be exposed bare, thereby giving a non-uniform zinc coating.

As to the Item 4, when the zinc powders adhere on the heated steel sheet, the iron atoms and the zinc atoms diffuse mutually to form an alloy layer. If this is to be prevented, it is known that the formation of Zn-Al compounds or the like on the boundary between the coated layer and the steel sheet is effective. If this is to be ensured, the aluminum content within the zinc powders should be preferably limited to 0.1 - 0.7 wt%.

If the Al content is less than 0.1%, an alloy layer is developed on the boundary, and therefore, the close adherence in the zinc coating is aggravated. On the other hand, if the Al content is more than 0.7%, although there is no problem in the formation of the coated layer, the formed coated layer is a Zn-Al alloy layer rather than a pure zinc layer. Therefore, the coated sheet is not suitable for use in automobiles and electronic apparatuses.

If the Item 5 is to be met, the steel sheet should be subjected to a reheating after its passage through the zinc coating chamber. If this step is omitted, the imperfectly adhered zinc powders can be transferred to various rolls to cause defects such as dent or the like. As to the reheating conditions, the reheating is carried out at a temperature of 420 - 650°C for 1 - 20 seconds. More precise conditions are

decided by the composition of the target coated layer. That is, if the target coated layer is a pure zinc layer, then the steel sheet may be heated to 420 - 500°C. Then the zinc powders are completely melt-adhered to form an acceptable zinc coated layer. When a Zn-Fe alloy coated layer is aimed at, the steel sheet is heated at 500 ~ 650°C for 10 ~ 20 seconds so as to promote the alloying reactions. Thus the loosely adhered zinc powders are made to melt-adhere, as well as promoting the alloying reactions. In this way, the reheating is carried out suitably with the composition of the coated layer after the steel sheet has passed through the zinc coating chamber as described above. Therefore, the loosely adhered zinc powders are converted into a coated layer, and at the same time, a coated layer having the target Fe content can be obtained.

In the present invention, the interior of the zinc coating chamber has to be filled with an inert gas or a reducing gas, and has to be filled with fluidized zinc powders. The temperature of the zinc coating chamber should be preferably limited to below 250°C.

If the zinc powders are contacted to the steel sheet based on the general method such as spouting or the like, locally non-uniform portions will necessarily occur due to the differences in the flow pattern, thereby making it difficult to obtain a uniform coated layer. Therefore, the present inventors studied on the method of contacting the zinc powders to the steel sheet, and as a result, found the following fact. That is, if fluidized zinc powders like a fog are uniformly

dispersed within the zinc coating chamber, and if the steel sheet is made to pass through the fog, then a uniform coated layer could be obtained.

Under this condition, the gas which is used for forming the fluidized bed of the zinc powders should be a reducing gas or a non-oxidizing gas. Otherwise, oxidation reactions occur on the surfaces of the steel sheet, and consequently, the adherence of the coated layer is aggravated. Further, if the temperature of the fluidized bed of the zinc powders exceeds 250°C, then the fluidized zinc powders are liable to be agglomerated as shown in FIG. 6, with the result that the stable fluidizing is destroyed. Consequently, the zinc particles adhere on the manufacturing facility in the form of agglomerates.

After satisfying the Items 1, 2, 3, 4 and 5, if the zinc coating method is to have an efficiency, the desired coated amount has to be obtained in an easy manner.

In the present invention, if the above conditions are all met, it was confirmed as shown in FIG. 7 that the coated amount can be adjusted by adjusting the flow rate of the gas which is used for forming the fluidized zinc bed. FIG. 7 is a graphical illustration showing the variation of the coated amount versus the variation of the gas flow rate, when the temperatures of the coated steel sheets are different. This is an evidence to the fact that, if all the conditions of the present invention are satisfied, as the gas flow rate increases, the movements of the zinc powders become brisk, and the zinc particles

collide with the steel sheet in an increased amount. That is, if the temperature of the steel sheet is below  $420\,^{\circ}\text{C}$ , the diffusion velocities become slow, and therefore, the coated speed becomes slow. Therefore, a coated amount of  $100~\text{g/m}^2$  or more within 5 seconds cannot be obtained unlike the general cases.

Therefore, in the present invention, the temperature of the steel sheet is limited in view of the general zinc coated amount and the limit of the treatment time of the manufacturing facility.

Now the method for coating zinc on the steel sheet by utilizing the electrostatic attraction according to the present invention will be described.

If the zinc coating is to be carried out according to the present invention, zinc powders have to be supplied through the powder supply device 246, and the zinc powders have to be fluidized within the fluidized bed forming chamber 240 by using a gas which is spouted from below.

The reason why the zinc powders have to be fluidized in advance is as follows.

The zinc powders have a naturally agglomerating trend, and therefore, if they are spouted as they are stored, then they are agglomerated into large particles so as to form coarse secondary particles. If such coarse secondary particles are spouted, the electrostatic attractions cannot give a satisfactory effect. Further, coating differences are generated over the different parts of the steel sheet, and

therefore, a uniform coated layer can hardly be obtained.

Therefore, the present inventors studied on the method of carrying the zinc powders. As a result of the study, the present inventors found the following facts. That is, if the fluidized bed forming technique is employed, then particles of more than a certain size can be prevented from entering into the zinc coating chamber. Therefore, in the present invention, the separate fluidized bed forming chamber 240 is provided separately from the zinc coating chamber 220. Then the zinc powders are carried from the fluidized bed forming chamber 240 to the zinc coating chamber 220.

The size of the particles suspended within the fluidized bed forming chamber 240 is closely related to the pressure of the gas which is spouted from below. As the pressure of the spouted gas increases, so the size of the suspended particles increases.

Therefore, if the pressure of the spouted gas is adjusted, then the coarse secondary particles can be made to sink onto the bottom, and the particles of the desired size can be suspended, so that they can be carried to the zinc coating chamber. Further, an injection pump 223b which is based on the principle of the ventury tube is installed on the zinc powder carrying path between the fluidized bed forming chamber 240 and the zinc coating chamber 220. Then the agglomerates receive mechanical impacts from an auxiliary gas so as to be disintegrated into individual particles. Therefore, the survival opportunity of the coarse secondary particles is further diminished, thereby

giving a uniform distribution of the particles during the carriage of the zinc powders. Thus only fine primary zinc particles adhere on the steel sheet to be converted into a coated layer. Then the microstructure of the coated layer becomes more uniform, and the melting speed of the zinc powders becomes faster. Then the departure of the zinc particles and therefore, the method becomes more becomes rarer, advantageous for a thick zinc coating. The external appearance of the coated steel sheet is also improved by the uniform suspension of the zinc powders within the gas. Further, electrostatic effect on the particles becomes greater, therefore, the influence of the spouting track decreases, with the result that a more uniform coated layer can be obtained.

As described above, the zinc powders which have been fluidized in the fluidized bed forming chamber 240 are spouted into the zinc coating chamber 220 through the powder spouting device 223. Thus the zinc powders maintain a suspended state within the zinc coating chamber 220, and the zinc powders are electrostatically charged.

For this purpose, the temperature of the steel sheet is maintained above the melting point of zinc, and the zinc coating power is the force of the carrying gas and the electrostatic attractions.

Owing to a temperature difference between the zinc powders and the steel sheet, a convection boundary layer is formed on the surfaces of the steel sheet, and the mentioned layer obstructs the access of the zinc powders. Therefore, in a

method in which the powders are spouted in a simple manner, the pressure of the carrier gas has to be increased, so that the powders can overcome the boundary layer to adhere on the steel sheet. In this case, differences in the spouting tracks occur over different parts of the steel sheet, and consequently, the coated layer becomes non-uniform. However, the electrostatic attraction is proportional to the square of the distance between two charged objects (F  $\infty$  1/r<sup>2</sup>). Therefore, the electrostatic attractive force is large near the convection boundary layer of the steel sheet, and therefore, the zinc powders which are carried to the convection boundary layer by the carrier gas easily adhere on the steel sheet. If this electrostatic attraction is utilized, the zinc powders adhere on the steel sheet even if the pressure of the carrier gas is lowered to as low as not to affect the steel sheet, and the spouting tracks do not appear on the surface of the steel sheet. powders which are sequentially adhere on the steel sheet meltadhere on the steel sheet in a sequential manner before the powder lose the electrostatic charges. Thus the zinc powders are firmly melt-attached on the steel sheet, and therefore, detachments of the zinc particles due to the dissipation of the electrostatic charges do not occur. Therefore, the method of the present invention is advantageous for a thick zinc coating.

In the present invention, the zinc powders are fluidized, and the zinc powders are electrostatically charged as described above. Therefore, the zinc coating can be done more speedily compared with the conventional method.

Then the steel sheet is heated to 420 ~ 730°C and grounded. This steel sheet is made to pass through the electrostatically charged fluidized zinc powders, so that the zinc powders would be melt-attached so as to form a coated layer.

Meanwhile, the residual uncoated zinc powders which remain on the bottom are sent to the cyclone. In the cyclone, the zinc powders are separated from the gas, and the gas is discharged, while the separated zinc powders are sent to the powder supply device 246, thereby recovering the zinc powders.

In the present invention, the zinc powders should be preferably limited to an average size of 45  $\mu m$ . The average size of 45  $\mu m$  cannot be applied to a small coated-amount zinc coating, because in this case some parts of the steel sheet may be exposed.

Further, when the zinc powders melt-adhere, iron and zinc atoms mutually diffuse to form an alloy layer, and therefore, this phenomenon needs to be inhibited. Thus Fe-Al or Fe-Al-Zn compounds may be formed on the boundary between the coated layer and the steel sheet.

The influence of an aluminum content on the zinc powders was studied, and the result showed the following fact. That is, the Al content should be preferably limited to 0.1-0.7 wt%. There is no problem in forming the coated layer, even if the Al content is more than 0.7%. However, in this case, the coated layer is not a zinc coated layer, but an Al-Zn alloy coated layer.

In a zinc coating, the zinc particles adhered on the steel

sheet are transferred to various rolls to cause defects such as dent. If the coated steel sheet is reheated, then the defects such as dent can be avoided. According to experiments, if the reheating conditions are precisely adjusted, then the composition of the coated layer can be varied. That is, in the case where a pure zinc coating is aimed at, the steel sheet is heated at 420-500°C for 1-5 seconds, and then, is cooled. Then only the zinc powders can be coated without inviting the alloying reactions. On the other hand, in the case where a Zn-Fe alloy coated layer is the target, the coated steel sheet is heated at 500-650°C for 10 to 20 seconds so as to promote the alloying reactions.

The heating period of the pure zinc coating is shorter than that of an alloy zinc coating, and therefore, the attachment of the zinc particles on the rolls becomes more probable. However, if a wash is carried out before the steel sheet contact with the rolls, then the loosely attached zinc powders can be completely removed. Therefore, the conventional problems such as the attachment of the zinc particles on the deflector roll or the peeling of the coated layer can be completely solved.

In the present invention, the internal atmosphere of the zinc coating chamber consists of an inert gas or a reducing gas, while internal temperature of the zinc coating chamber is the normal temperature to 250°C. The reason is as follows. That is, if the temperature of the steel sheet during the zinc coating drops to below the melting point of zinc, then the

adhering efficiency of the zinc powders decreases. In order to prevent this, the atmospheric gas should have a temperature as high as possible. However, if the temperature of the atmospheric gas exceeds 250°C, the fluidized zinc particles tend to be agglomerated as shown in FIG. 6, thereby aggravating the stability of the fluidized bed. The optimum temperature of the atmospheric gas is 100-200°C.

Under the above conditions, the flow rate of the gas and the applied voltage of the electrodes are adjusted to adjust the coated amount.

FIG. 7 is a graphical illustration showing the variation of the coated amount versus the variation of the gas flow rate, when the fluidized zinc powders are injected into the zinc coating chamber. This drawing shows that as the gas flow rate increases, the adhered zinc amount increases, if the conditions of the present invention are satisfied.

FIG. 8 is a graphical illustration showing the variation of the coated amount versus the variation of the voltage of the electrode, in a state with the steel sheet grounded. As the applied voltage of the electrode increase, the zinc coated amount steeply increases, to such a degree that a coated amount of 200 g/m² can be easily obtained. Under this condition, in electrostatically charging the zinc powders, the corona charging or the induction charging is employed. For this purpose, a sharp tipped nozzle and a net type electrode are used. As the applied voltage of the electrode, -1 ~ -100 KV or 1 ~ 100 KV will be enough.

When the zinc powders are used as in the case of the present invention, the zinc powders can be introduced into the annealing furnace. Consequently, the zinc powders can adhere on various rolls to cause defects such as dent. Further, if the zinc powders are leaked to the outside of the manufacturing facility, the powders may hurt the health of workers. Therefore, the recovery of the zinc powders is very important. In this context, the internal pressure of the zinc coating chamber should be properly adjusted, and the leakage of the zinc powders should be prevented. That is, measures for these should be prepared. Accordingly in the present invention, the sealing chambers are provided above and below the zinc coating chamber, and the zinc powder recovering device is installed.

Now the present invention will be described based on actual example.

### <Example 1>

A cold rolled steel strip was heated to the temperatures of Tables 1 and 2. Then it was passed through a fluidized bed of zinc powders to coat it up to the optimum coated amount. Then the coated steel strip was reheated, thereby preparing coated test pieces. In varying the coated amount, the relationship between the coated amount and the gas flow rate as shown in FIG. 6 was utilized.

Tables 3 and 4 shows the effects of the zinc coating conditions.

The adherence strength of the coated layer was evaluated

based on a 45-degree bending test, i.e. based on the peeling degree during the bending test. The level of the absolutely non-peeling of the coated layer was shown by " $\odot$ ". The level in which the traces of peeling appeared was shown by "O". The level in which the traces of peeling definitely appeared was shown by " $\Delta$ ". The level in which the coated layer was almost peeled off was shown by "X".

As to the coating uniformness, the external appearance was observed by human eyes, and the structure of the coated layer was observed by magnifying it to 2000 times by a scanning microscope. Thus, if it has a uniform structure without any pin hole, then it was assigned with "o". If the external appearance was uniform, but if the structure was not uniform, then it was assigned with "O". If both the external appearance and the structure were not uniform, it was assigned with "A". If a coated layer was not formed at all, the it was assigned with "X".

The coatability indicates the maximum coated amount which can be obtained within 5 seconds as allowed in the general continuous annealing factory. "X" indicates the case where a coated layer was not formed at all. " $\Delta$ " indicates the case where a thin coated layer of less than 40 g/m² was obtained. "O" indicates the case where the desired coated amount was obtained by varying the zinc coating conditions.

The paintability was evaluated in such a manner that a melamine alkydic pigment was spread, and then, straight scratchings were carried out at intervals of 1 mm in the form

of check works. Then the evaluation was carried out.

< Table 1 >

				Coated	layer form	ning condit	ions	
		Sheet tempera- ture (°C)	Size of Zn powder (μm)	content	Atmos- phere	Temper- ature of fluidized bed	Heating time (sec )	Reheating temper- ature (°C)
	1	740	5	<0.01	N <sub>2</sub> ·	200	non-rel	neating
	2	740	5	0.18	N2+H2	200	non-rel	
	3	500	5	0.18	N2+H2	100	1	410
	4	500	5	<0.01	N <sub>2</sub>	100	5	410
	5	420	40	<0.01	N <sub>2</sub>	100	25	410
	6	500	40	0.18	N <sub>2</sub>	100	25	650
	7	500	0.5	0.14	N <sub>2</sub>	100	25	410
	8	390	20	0.18	N <sub>2</sub>	100	5	520
a	9	390	5	0.18	N <sub>2</sub>	100	10	520
աթյ	10	740	5	0.18	N <sub>2</sub>	150	5	520
example	11	740	5	0.18	N <sub>2</sub>	100	10	520
o V	12	450	5	<0.01	N <sub>2</sub>	100	5	520
ati,	13	600	20	0.18	N <sub>2</sub>	100	5	660
Comparative	14	450	20	0.18	N2+H2	200	25	420
dino	15	500	50	0.14	N2+H2	200	15	550
O	16	550	50	0.18	N2+H2	200	15	520
	17	730	70	0.18	N2+H2	200	2	650
	18	500	20	<0.01	N2+H2	200	- 15	520
	19	500	5	<0.01	N2+H2	200	12	520
	20	500	5	0.18	oxidizing	200	15	520
	21	730	5	0.18	oxidizing	200	8	650
	22	730	5	0.18	N2+H2	300	8	650
	23	550	5	0.14	N2+H2	300	15	600

< Table 2 >

		Coated layer forming conditions						
		Sheet	Size	Al	Atmos-	Temper-	Heating	Reheating
		temper-	of Zn	content	phere	ture of	time	temper-
	ļ	ature	powder	within Zn		fluidized	(sec )	ature
		(℃)	$(\mu_m)$	powder		bed		(℃)
		500		(wt.%)				
	1	500	5	0.18	N <sub>2</sub> +H <sub>2</sub>	100	1	420
	2	500	5	0.1	N <sub>2</sub>	100	5	420
	3	420	40	0.1	N <sub>2</sub>	100	20	420
	4	500	40	0.18	N <sub>2</sub>	100	20	650
	5	500	0.5	0.14	N <sub>2</sub>	100	20	420
	6	420	20	0.18	N <sub>2</sub>	100	5	520
	7	420	5	0.18	N <sub>2</sub>	100	10	520
	8	730	5	0.18	N <sub>2</sub>	150	5	520
ple	9	730	5	0.18	N <sub>2</sub>	100	10	520
example	10	450	5	0.1	N <sub>2</sub>	100	5	520
1	11	450	20	0.18	N <sub>2</sub>	100	5	420
Inventive	12	600	20	0.18	N <sub>2</sub>	100	5	650
nve	13	450	20	0.18	N <sub>2</sub>	200	5	420
	14	450	20	0.18	N <sub>2</sub>	200	5	600
	15	450	20	0.18	N2+H2	200	20	420
	16	500	0.5	0.14	N2+H2	200	15	520
	17	500	45	0.14	N2+H2	200	15	550
	18	730	45	0.18	N2+H2	200	2	650
	19	500	20	0.1	N2+H2	200	15	520
	20	730	5	0.7	N2+H2	200	15	650
	21	450	5	0.7	N2+H2	200	20	500

< Table 3 >

		Adherence of coated layer	Uniformness of coated layer	Coatability	Paintability
	1	Δ	0	0 '	Δ
	2	Δ	O.	0	Δ
	3	Δ .	0	0	Δ
	4	Δ	0	0	Δ
	5	×	0	0	Δ
	6	Δ	0	0	0
	7	, Δ .	0	0	0
	8	0	Ο.	Δ	0
ple	9	0	0	Δ	0
example	10	. 🛆	Δ	0	0
	11	_ Δ	Δ	0	0
Comparative	12	Δ	0	0	0
para	13	Δ	0	Δ	0
Com	14	Δ	0	0	0
	15	0	×	0	0
	16	· ©	×	0	0
	17	0	×	Ó	0
	18	×	0	0	0
	19	×	0	0	0
	20	×	×	×	×
	21	×	. ×	×	×
	22	0	0	0	0
	23	0	0	0	0

 $\bigcirc$ : Excellent,  $\bigcirc$ : Adequate,  $\triangle$ : Bad, X: Very bed

< Table 4 >

		Adherence of coated layer	Uniformness of coated layer	Coatability	Paintability
	1	0	0	0	0
	2	0	0	0 .	0
	3	0	0	0	0
	4	0	0	0	0
	5	. ©	0	0	0
	6	0	0	0	0
	7	<b>©</b> .	0		. ©
}	8	0	0	0	· ©
] e	9	©	0	0	0
example	10	0	0	0	0
1	11	©	0	0	0
ntiv	12	© .	0	0	0
Inventive	13	0	0	0	0
	14	0	0	0	0
	15	0	0	0	0
	16	0	0	0	0
	17	0	0	0 .	0
	18	0	0	0	0
	19	· ©	0	0	0
	20	0	0	0	0
	21	0	© .	0	0

 $\bigcirc$ : Excellent,  $\bigcirc$ : Adequate,  $\triangle$ : Bad, X: Yery bed

Table 3 shows the results of evaluations of the qualities of the coated layer for Comparative examples 1-23 which were manufactured at the conditions of Table 1. As shown in Table 3, at least one among the adherence of the coated layer, the uniformness of the coated layer, the coatability and the paintability was defective. This is due to the fact that at least one item among them departed from the range of the present invention. On the other hand, the coated layers which were manufactured based on the method of the present invention as shown in Table 2 were all satisfactory as shown in Table 4 in all the respects including the adherence of the coated layer, the uniformness of the coated layer, the coatability and the paintability.

### <Example 2>

A zinc coating was carried out at conditions same as those of the inventive example 1 of Table 2, except that the sheet temperature and the gas flow rate were varied as shown in FIG. 6. The variation of the coated amount versus the variation of the gas flow rate for the fluidized bed was checked, and the results are shown in FIG. 6.

As shown in FIG. 6, if the method of the present invention is applied, the coated amount increases as the gas flow rate increases.

### <Example 3>

A zinc coating was carried out by using the zinc coating apparatus of FIG. 4 at the conditions of Tables 5 and 6. Then the adherence of the coated layer, the uniformness of the coated layer, the coated layer, the paintability were evaluated, and the results are shown in Tables 7 and 8 below.

In tables 5 and 6, the comparative examples and the inventive examples 22-29 used an electrode voltage of -55 KV, while the inventive examples 30-31 used an electrode voltage of -90 KV.

Meanwhile, the flow rate of the fluidized bed forming gas was 100 L/min, while the flow rate of the auxiliary gas at the injection pump was 100 L/min.

< Table 5 >

		Coated layer forming conditions						
		Sheet temper- ature (°C)	Size of Zn powder (μm)	Al content within Zn powder (wt.%)	Atmos- phere	Temper- ature of fluidized bed	Heating time (sec )	Reheating temper- ature (°C)
	24	740	5	0.18	N <sub>2</sub>	100	non-re	eheating
به	25	410	5	0.18	N <sub>2</sub>	100	25	550
example	26	550	50	0.18	N <sub>2</sub>	150	1	. 550
	27	550	15	0.07	N <sub>2</sub>	200	non-re	heating
tive	28	720	15	0.8	N <sub>2</sub>	100 .	5	550
para	29	600	15	0.14	oxidizir	g 100	10	550
Compara	30	600	5	0.5	N <sub>2</sub>	250	5	600
	31	550	5	0.3	N <sub>2</sub>	100	25	650

# < Table 6 >

			ons					
		Sheet	Size of	Al	Atmos-	Temper-	Heating	Reheat ing
		temper-	Zn	content	phere	ature of	time	temper-
		ature	powder	within	•	fluidized	(sec )	ature
		(℃)	(μm)	Zn powder	-	bed (℃)	-	(℃)
				(wt.%)		<u>'</u>		
	22	550	5	0.18	N <sub>2</sub>	. 120	5	550
	23	720	5	0.18	N <sub>2</sub>	100	non-reheating	
	24	420	5	0.18	N <sub>2</sub>	100	non-reheating	
)le	25	550	15	0.18	N <sub>2</sub>	150	10	550
example	26	550	15	0.1	N2 -	200	1	650
i :	27	550	15	0.7	N <sub>2</sub>	100	5	550
Inventive	28	600	45	0.14	N <sub>2</sub>	100	10	550
Inve	29	600	5	0.5	N <sub>2</sub>	200	5	600
	30	550	5	0.3	N2	100	20	650
	31	550	5	0.2	N <sub>2</sub>	100	non-r	eheating

## < Table 7 >

		Adherence of coated layer	Uniformness of coated layer	Coated amount (g/m')	Paintability
	24	Δ	0	200	0
nple	25	0	Δ .	80	0
exampl	26	0	Δ	220	Δ
o V	27	×	0	200	0
rati	28	0	0	200	×
Compar	29	×	×	-	×
ß	30	0	Δ	200	. 0
	31	Δ	0 .	200	Δ

(7) (1) (1)

# < Table 8 >

		Adherence of coated layer	Uniformness of coated layer	Coated amount (g/m')	Paintability
	22	©	0	200	   ©
	23	0	0	200	0
ple	24	0	0	220	0
example	25	0	0	200	0
ve e	26	0	<u>.</u> ©	200	0
1:	27	<b>©</b>	0	200	0
Inven	28	0	. 0	200	0
I	29	0	0	200	0
	30	© .	0	300	0
	31	0	0	300	0

As shown in Table 7 above, the coated amount is about 200  $g/m^2$  in all of them, except the comparative example 25 which shows a low coated amount.

The comparative example 25 shows a coated amount as low as  $80~\text{g/m}^2$ , and this is due to the fact that the adhered zinc powders are detached before they are converted into a coated layer.

Meanwhile, the comparative examples 24-31 show one or more defects among the adherence of the coated layer, the uniformness of the coated layer, and the paintability. This is due to the fact that they departed from at least one or more of the coated layer forming conditions of the present invention.

On the other hand, as shown in Table 8, the inventive examples achieved the coated layers of more than  $200~\text{g/m}^2$  in all of them. Particularly, in the inventive examples 30 and 31 in which a voltage of -90~KV was applied, a coated amount of  $300~\text{g/m}^2$  was obtained.

In the cases of the inventive examples 22, 23 and 31 in which the reheating was not carried out, some individual zinc particles were observed, but the adherence of the coated layer and the paintability were all satisfactory.

In the cases of the other inventive examples, the adherence of the coated layer, the uniformness of the coated layer, and the paintability were all satisfactory.

### <Example 4>

A zinc coating was carried out at conditions same as those

of the inventive example 22 of Table 6, except that the gas flow rate and the electrode voltage were varied as shown in FIG. 8. The evaluated results are shown in FIG. 8.

As shown in FIG. 8, in the present invention, a coated amount of 200  $g/m^2$  or more could be easily obtained.

According to the present invention as described above, a zinc coating apparatus and a method therefor are provided in which the zinc coating speed is as fast as to be connected a continuous annealing furnace for cold rolled steel sheet, the coating deviations are smaller than those of the hot dip galvanizing apparatus, and a thick coated layer can be easily formed. Therefore, compared with the conventional method, the present invention improves the product quality and the productivity.

#### WHAT IS CLAIMED IS:

 An apparatus for continuously coating zinc on a steel sheet, comprising:

a zinc coating chamber for forming a fluidized bed of zinc powders, for passing a heated steel sheet (heated by a heating means) through the fluidized bed of the zinc powders, and for making the zinc powders melt-adhere on the steel sheet during a passing of the steel sheet through the fluidized bed;

a fluidized bed forming chamber for forming a fluidized bed of the zinc powders by making the zinc powders suspended by spouting a gas;

a cyclone for separating the zinc powders from the gas after recovery of them from said zinc coating chamber, to discharge the gas, and to return the separated zinc powders to said fluidized bed forming chamber;

a deflector roll for shifting an advancing direction of the steel sheet after its admittance into said zinc coating chamber;

a tension roll for shifting an advancing direction of a zinc coated steel sheet;

said zinc coating chamber comprising: a powder inlet tube connected from a side wall of said zinc coating chamber to the fluidized bed to inject the zinc powders into said zinc coating chamber; a gas inlet tube for forming a turbulent flow of the zinc powders and for preventing a leakage of the zinc powders; and a recovering tube for recovering uncoated zinc powders;

said gas inlet tube being disposed above said powder inlet tube, and said recovering tube being disposed below said powder

inlet tube;

said recovering tube being connected between said zinc coating chamber and said cyclone, and a suction pump being connected to said cyclone;

a separating plate provided within said zinc coating chamber, for making uncoated zinc powders smoothly flow to said recovering tube, and for preventing the zinc powders from flowing to a stabilizing roll; and

a stabilizing roll disposed below said separating plate.

- 2. The apparatus as claimed in claim 1, wherein said heating means is an annealing furnace.
- 3. The apparatus as claimed in any one of claims 1 and 2, wherein a reheating furnace is installed between said zinc coating chamber and said tension roll.
- 4. An apparatus for continuously coating zinc on a steel sheet, comprising:

a zinc coating chamber for making zinc powders melt-adhere on a heated steel sheet to form a coated layer;

a fluidized bed forming chamber for forming a fluidized bed of the zinc powders by making the zinc powders suspended by spouting a gas;

a cyclone for separating the zinc powders from the gas after recovery of them from said zinc coating chamber, to discharge the gas, and to return the separated zinc powders to

said fluidized bed forming chamber;

a deflector roll for shifting an advancing direction of the steel sheet after its admittance into said zinc coating chamber;

a tension roll for shifting an advancing direction of a zinc coated steel sheet;

said zinc coating chamber comprising a powder spouting tube connected from a side wall of said zinc coating chamber to the fluidized bed to spout the zinc powders into said zinc coating chamber;

said zinc coating chamber further comprising a recovering tube connected to said cyclone, for recovering uncoated zinc powders; and

one or more of electrodes provided in said zinc coating chamber, for electrostatically charging the zinc powders, said electrodes being connected to a high voltage generating device.

- 5. The apparatus as claimed in claim 4, wherein a reheating furnace is installed between said zinc coating chamber and said tension roll.
- 6. The apparatus as claimed in any one of claims 4 and 5, wherein said powder spouting device comprises: a powder carrying tube connected to said fluidized bed forming chamber; an injection pump connected to said powder carrying tube; and a powder spouting nozzle for spouting the zinc powders from said injection pump into said zinc coating chamber.

- 7. The apparatus as claimed in any one of claims 4 and 5, wherein said electrode is installed within said zinc coating chamber, in a state insulated from said zinc coating chamber a pair of sharp electrodes of a side wall of said zinc coating chamber.
- 8. The apparatus as claimed in claim 6, wherein said electrode consists of a pair of net type electrodes installed within said zinc coating chamber, in a state insulated from the pair of said sharp electrodes and from said zinc coating chamber.
- 9. The apparatus as claimed in claim 4, wherein a cooling device is installed beneath said zinc coating chamber.
- 10. The apparatus as claimed in claim 9, wherein said cooling device comprises: a water spouting nozzle; and an air spouting nozzle for forming an air curtain above said water spouting nozzle.
- 11. The apparatus as claimed in claim 5, wherein said cooling device is disposed below said reheating furnace.
- 12. The apparatus as claimed in claim 11, wherein said cooling device comprises: a water spouting nozzle; and an air spouting nozzle for forming an air curtain above said water spouting nozzle.

- 13. The apparatus as claimed in any one of claims 10 and 12, wherein a wash device is installed beneath said water spouting nozzle, for washing surfaces of the steel sheet.
- 14. A method for continuously coating zinc on a steel sheet by making the steel sheet pass through a zinc coating chamber, comprising the steps of:

supplying fluidized zinc powders of a fluidized bed forming chamber to said zinc coating chamber, and injecting an inert gas or a reducing gas into said zinc coating chamber through a side wall of said zinc coating chamber to form a fluidized bed within said zinc coating chamber;

making a heated steel sheet (heated to 420 - 730°C) through the fluidized bed within said zinc coating chamber to meltattach the zinc powders on the steel sheet so as to form a coated layer;

reheating the zinc powder-adhered steel sheet at a temperature of  $420-650\,^{\circ}\text{C}$  for 1-20 seconds to make residual uncoated zinc powders melt-adhere on the surface of the steel sheet so as to form a coated layer; and

discharging residual uncoated zinc powders from a bottom portion of said zinc coating chamber together with a gas by a cyclone, to separate the zinc powders from the gas so as to discharge the gas and so as to return the separated zinc powders to said fluidized bed forming chamber.

15. The method as claimed in claim 14, wherein the zinc

powders have an average particle size of 45  $\mu m$ .

- 16. The method as claimed in any one of claims 14 and 15, wherein the zinc powders contain Al in an amount of 0.1-0.7 wt%.
- 17. The method as claimed in any one of claims 14 and 15, wherein a reheating temperature of 420 500°C and a reheating time period of 1 5 seconds are adopted.
- 18. The method as claimed in claim 16, wherein a reheating temperature of 420 500°C and a reheating time period of 1 5 seconds are adopted.
- 19. The method as claimed in any one of claims 14 and 15, wherein a reheating temperature of 500 650°C and a reheating time period of 10 20 seconds are adopted.
- 20. The method as claimed in claim 16, wherein a reheating temperature of 500 650°C and a reheating time period of 10 20 seconds are adopted.
- 21. The method as claimed in any one of claims 14 and 15, wherein said zinc coating chamber adopts an atmospheric gas temperature of 250°C or below.
  - 22. The method as claimed in claim 16, wherein said zinc

coating chamber adopts an atmospheric gas temperature of 250°C or below.

23. A method for continuously coating zinc on a steel sheet by making the steel sheet pass through a zinc coating chamber, comprising the steps of:

receiving zinc powders from a powder supply device, and fluidizing the zinc powders within a fluidized bed forming chamber by a help of a gas blown through a lower portion of it;

injecting the fluidized zinc powders from said fluidized bed forming chamber to a zinc coating chamber by means of an injecting device to form a fluidized bed within said zinc coating chamber;

charging the zinc powders of the fluidized bed positively or negatively;

heating a steel sheet to 420 - 730°C and grounding the steel sheet, and making the steel sheet pass through the fluidized bed to make the charged zinc powders melt-adhere on the steel sheet;

sending a residual zinc powders of a bottom portion of said zinc coating chamber to a cyclone together with a gas to separate the zinc powders from the gas, so as to discharge the gas, and so as to return the separated zinc powders to a powder supply device.

24. The method as claimed in claim 23, wherein the zinc powders have an average particle size of 45  $\mu m$ .

- 25. The method as claimed in any one of claims 23 and 24, wherein the zinc powders contain Al in an amount of 0.1 0.7 wt%.
- 26. The method as claimed in claim 23, wherein after zinc-coating the steel sheet, the steel sheet is reheated at a temperature of 420 650°C, for making loosely attached uncoated zinc powders melt-adhere on the steel sheet.
- 27. The method as claimed in claim 26, wherein a reheating temperature of 420 500 °C and a reheating time period of 1 5 seconds are adopted.
- 28. The method as claimed in claim 26, wherein a reheating temperature of  $500-650\,^{\circ}\text{C}$  and a reheating time period of 10-20 seconds are adopted.
- 29. The method as claimed in any one of claims 23 and 24, wherein the steel sheet is grounded, and said electrode is supplied with a voltage of -1 ~ -100 KV, or 1 ~ 100 KV.
- 30. The method as claimed in claims 25, wherein the steel sheet is grounded, and said electrode is supplied with a voltage of -1 ~ -100 KV, or 1 ~ 100 KV.
- 31. The method as claimed in any one of claims 26 to 28, wherein the steel sheet is grounded, and said electrode is

supplied with a voltage of -1  $^{\sim}$  -100 KV, or 1  $^{\sim}$  100 KV.

- 32. The method as claimed in any one of claims 23 and 24, wherein said zinc coating chamber adopts an atmospheric gas temperature of a normal temperature to 250°C.
- 33. The method as claimed in claim 25, wherein said zinc coating chamber adopts an atmospheric gas temperature of a normal temperature to 250°C.
- 34. The method as claimed in any one of claims 26 to 28, wherein said zinc coating chamber adopts an atmospheric gas temperature of a normal temperature to 250°C.
- 35. The method as claimed in any one of claims 23 and 24, wherein after being zinc-coated in said zinc coating chamber, the steel sheet is cooled by spouting water through a nozzle.
- 36. The method as claimed in claim 25, wherein after being zinc-coated in said zinc coating chamber, the steel sheet is cooled by spouting water through a nozzle.
- 37. The method as claimed in any one of claims 26 to 28, wherein after being reheated, the reheated steel sheet is cooled by spouting water through a nozzle.

このない はいかい はい はいない

38. The method as claimed in any one of claims 23 and 24,

wherein after being zinc-coated in said zinc coating chamber, the steel sheet is washed by water.

- 39. The method as claimed in claim 25, wherein after being zinc-coated in said zinc coating chamber, the steel sheet is washed by water.
- 40. The method as claimed in any one of claims 26 to 28, wherein after being reheated, the reheated steel sheet is washed by water.

## ABSTRACT OF THE DISCLOSURE

An apparatus and a method for coating zinc on steel sheets for use on automobiles and electronic apparatuses are disclosed. A fluidized bed forming chamber is provided to fluidize zinc powders in carrying out a zinc coating on a heated steel sheet, so that not only a uniformly coated layer but also a thick coated layer can be obtained. The apparatus for continuously coating zinc on a steel sheet according to the present invention includes a zinc coating chamber for forming a fluidized bed of zinc powders, for passing a heated steel sheet through the fluidized bed of the zinc powders, and for making the zinc powders melt-adhere on the steel sheet during its passing through the fluidized bed. A fluidized bed forming chamber forms a fluidized bed of the zinc powders by making the zinc powders suspended by spouting a gas. A cyclone separates the zinc powders from the gas after recovery of them from the zinc coating chamber, to discharge the gas, and to return the separated zinc powders to the fluidized bed forming chamber. A deflector shifts the advancing direction of the steel sheet after its admittance into the zinc coating chamber. A tension roll shifts the advancing direction of a zinc coated steel sheet. Further, the zinc coating chamber includes one or more electrodes for electrostatically charging the zinc powders.

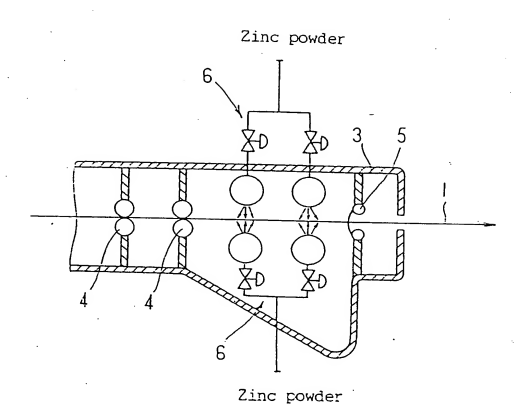


FIG.1

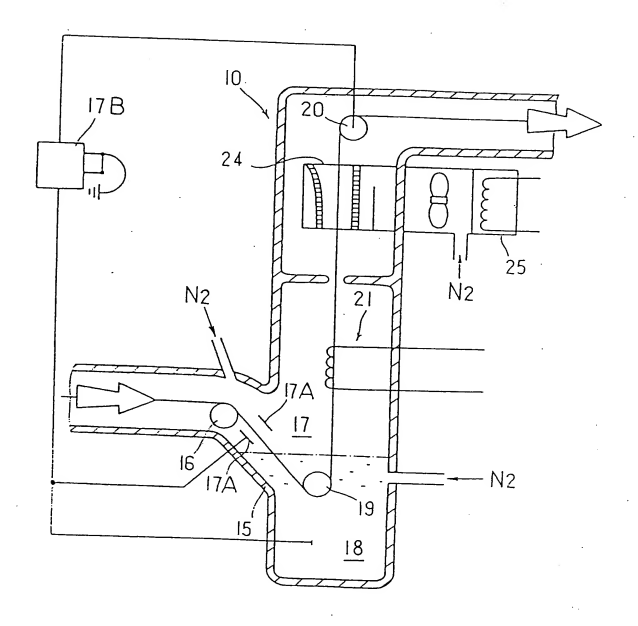
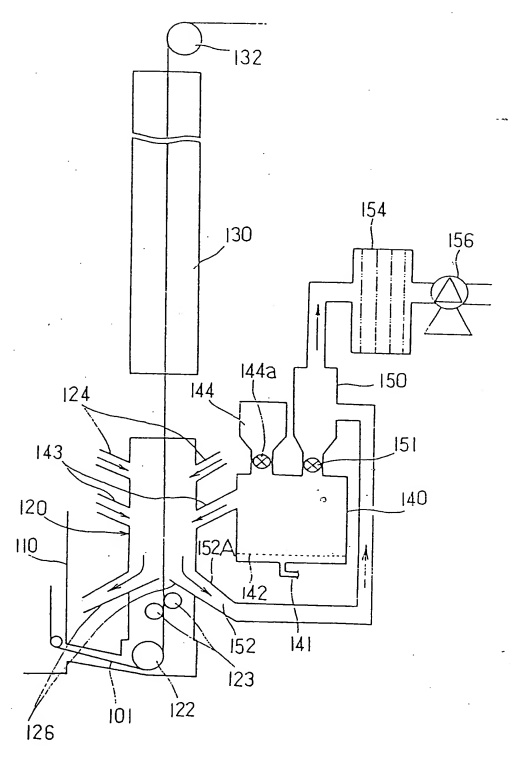
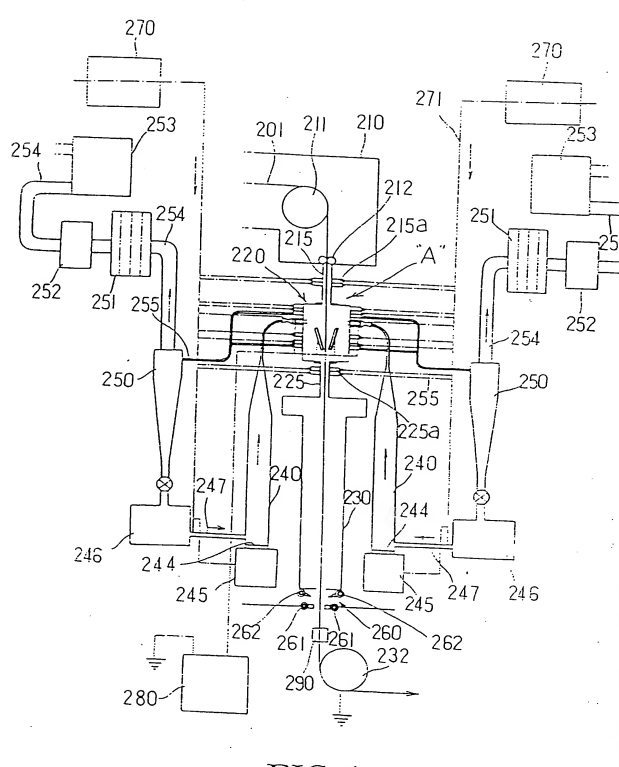


FIG.2



 $\binom{+}{2}$ 

FIG.3



 $\widehat{\left( \begin{array}{c} \cdot \cdot \cdot \\ \cdot \cdot \cdot \end{array} \right)}$ 

FIG.4

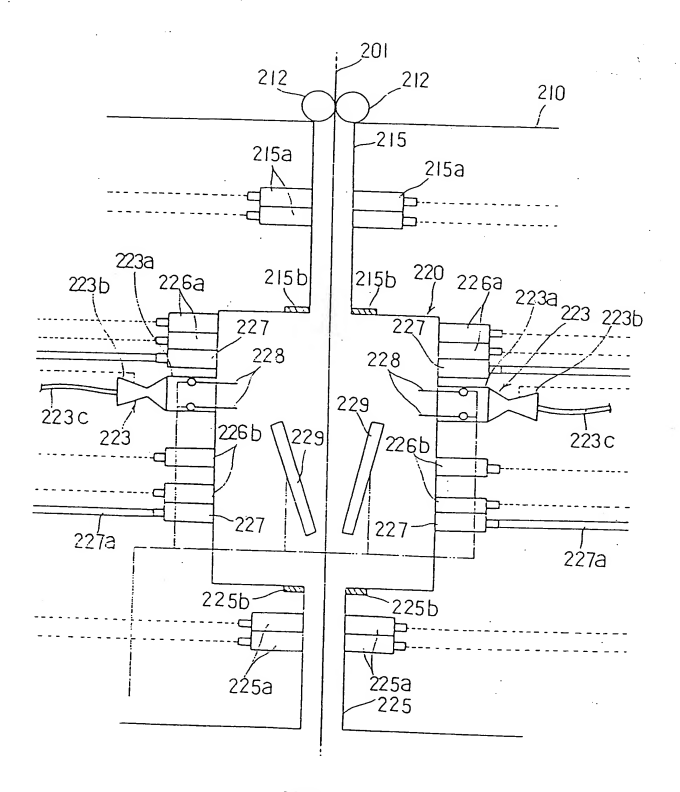
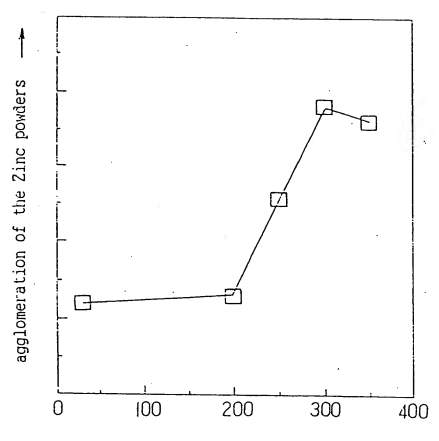


FIG.5



temperature of the fluidized bed (°C)

FIG.6

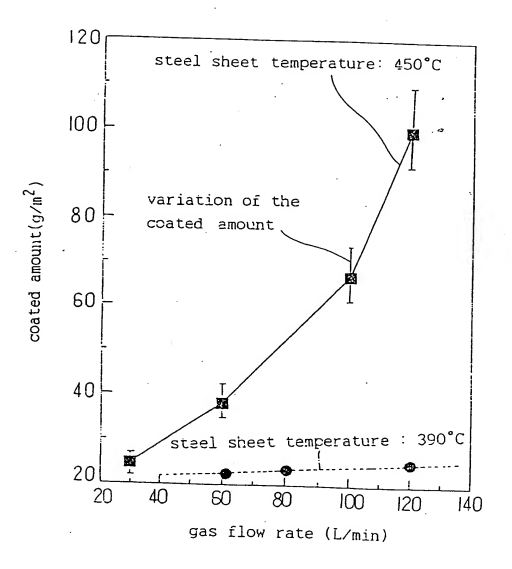


FIG.7

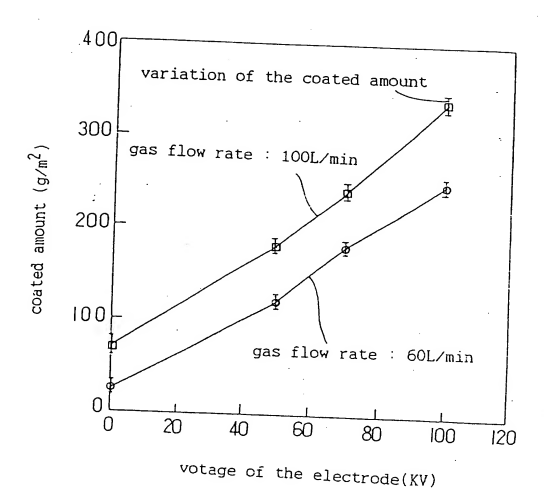


FIG.8